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OM nucleic - nucleic search, using sw model

Run on: March 9, 2002, 01:06:57 ; Search time 755.06 seconds

(without alignments)
28.386 Million cell updates/sec

Title: US-09-851-670-6

Perfect score: 25

Sequence: 1 cccctagccccaccagctctactgct 25

Scoring table: IDENTITY_NUC

Gapop 10.0 , Gapext 1.0

Searched: 930621 seqs, 428662619 residues

Total number of hits satisfying chosen parameters: 1026190

Minimum DB seq length: 0

Maximum DB seq length: 60

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : N.Geneseq.1101.*

1:	/SID2/gcgdata/geneseq/geneseqn/NA1980.DAT.*
2:	/SID2/gcgdata/geneseq/geneseqn/NA1981.DAT.*
3:	/SID2/gcgdata/geneseq/geneseqn/NA1982.DAT.*
4:	/SID2/gcgdata/geneseq/geneseqn/NA1983.DAT.*
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11:	/SID2/gcgdata/geneseq/geneseqn/NA1990.DAT.*
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13:	/SID2/gcgdata/geneseq/geneseqn/NA1992.DAT.*
14:	/SID2/gcgdata/geneseq/geneseqn/NA1993.DAT.*
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17:	/SID2/gcgdata/geneseq/geneseqn/NA1996.DAT.*
18:	/SID2/gcgdata/geneseq/geneseqn/NA1997.DAT.*
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20:	/SID2/gcgdata/geneseq/geneseqn/NA1999.DAT.*
21:	/SID2/gcgdata/geneseq/geneseqn/NA2000.DAT.*
22:	/SID2/gcgdata/geneseq/geneseqn/NA2001.DAT.*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query %	Match Length	ID	Description
1	18	72.0	18	AAF79646	Human Akt-3 antisense
2	16	64.0	25	AAAG6244	Human Akt-3 CDNA 3
3	15.2	60.8	20	AAO97960	PNA Oligomer target
4	15.2	60.8	20	AAO84237	PKC-epsilon coding
5	15.2	60.8	20	AAZ27354	Human protein kinase
6	15.2	60.8	20	AAZ78612	Human PKC-epsilon
7	15.2	60.8	20	AAZ83704	Human protein kinase
8	15.2	60.8	20	AAZ22650	Human protein kinase
9	15.2	60.8	20	AAZ19215	Human PKC-epsilon
10	15.2	60.8	36	AAT53201	Mouse ICM hammer
11	15.2	60.8	36	AAT52969	Mouse ICM hammer

C	12	15	60.0	32	22	AAH25021	PCR primer used to
C	13	15	59.2	51	22	AAH38540	Human SNP flanking
C	14	14.8	59.2	20	20	AAZ27368	Human protein kinase
C	15	14.8	59.2	20	20	AAZ78626	Human PKC-epsilon
C	16	14.8	59.2	20	20	AAZ83747	Human protein kinase
C	17	14.8	59.2	20	20	AAZ22664	Human protein kinase
C	18	14.8	59.2	20	20	AAZ19229	Human PKC-epsilon
C	19	14.6	58.4	38	17	AAZ64355	Rabbit streptolysin
C	20	14.4	57.6	51	15	AAO67302	PCR primer for pre
C	21	14.2	56.8	36	16	AAT54382	Human IL-5 hamster
C	22	14.2	56.8	54	15	AAO67301	PCR primer for pre
C	23	14	56.0	18	22	AAZ79647	Human Akt-3 antisense
C	24	14	56.0	26	22	AAZ17379	Information carboxyl
C	25	14	56.0	28	18	AAT76384	Human tumour necro
C	26	14	56.0	28	20	AAZ54533	Tumour necrosis fa
C	27	14	56.0	28	21	AAZ20099	Human tumour necro
C	28	14	56.0	28	21	AAZ33977	Low adenosine anti
C	29	14	56.0	51	22	AAH89369	Human nucleoside cod
C	30	14	56.0	57	15	AAO67295	PCR primer HCDRD5
C	31	14	56.0	57	15	AAO70503	Mutagenic primer p
C	32	14	56.0	57	17	AAT48542	Primer used for pr
C	33	13.8	55.2	18	20	AAZ18204	Serine threonine k
C	34	13.8	55.2	27	22	AAH38539	SNP specific SNPE
C	35	13.6	54.4	20	19	AAZ21330	Chimeric Ig germli
C	36	13.6	54.4	24	22	AAZ1892	Monoclonal anti
C	37	13.6	54.4	35	19	AAZ54220	Primer KC101 used
C	38	13.6	54.4	35	20	AAZ53502	Soluble SC-PCR fus
C	39	13.6	54.4	36	16	AAT55434	Human re1A hamster
C	40	13.6	54.4	36	16	AAT53290	Mouse ICM hammer
C	41	13.6	54.4	36	17	AAT52939	Human ICM hammer
C	42	13.6	54.4	36	17	AAZ65918	Human B7-2 hamster
C	43	13.6	54.4	36	17	AAZ65032	Human B7-1 hamster
C	44	13.6	54.4	36	17	AAT30349	Human YAP WW domai
C	45	13.6	54.4	38	17	AAZ64090	Rabbit stromelysin

ALIGNMENTS

RESULT 1
AAF79646 standard; DNA: 18 BP.

AC	AAF79646;
XX	
DT	29-MAY-2001 (first entry)
XX	
DE	Human Akt-3 antisense oligonucleotide, SEQ ID NO: 54.
XX	
KW	Human; Akt-3; protein kinase; cytosolic; antinflammatory; infection;
KW	antisense therapy; inflammation; tumour; ss.
XX	
OS	Homo sapiens.
XX	
PN	US6187585-B1
XX	
PD	13-FEB-2001.
XX	
PR	29-DEC-1999; 99US-0474922.
XX	
PR	29-DEC-1999; 96US-0474922.
XX	
XX	(ISIS-) ISIS PHARM INC.
XX	
PI	Monia BP, Cowsett LM, Roth RA;
XX	
DR	WPI: 2001-264979/27.
XX	
PT	New antisense compounds targeting nucleic acids encoding human Akt-3
PT	useful for treating a disease or condition associated with Akt-3
PT	expression, or in preventing or delaying inflammation or tumor
PT	formation

PS claim 1; Column 39; 37pp; English.

XX The present sequence is one of a number of antisense compounds of up to
CC 30 nucleobases in length targeted to a nucleic acid encoding human Akt-3.
CC The antisense compounds are useful for inhibiting the expression of human
CC Akt-3 in human cells or tissues. They are also useful for modulating the
CC expression of Akt-3, and for treating a human or an animal suspected of
CC having, or being prone to, a disease or condition associated with Akt-3
CC expression. The antisense compounds may also be used as research
CC reagents, in kits and in diagnostics, e.g. to elucidate the function of a
CC particular gene or to distinguish between functions of various members of
CC a biological pathway, and as a prophylactic, e.g. to prevent or delay
CC infection, inflammation or tumour formation.

XX Sequence 18 BP; 4 A; 8 C; 3 G; 3 T; 0 other;

XX Query Match 72.0%; Score 18; DB 22; Length 18;
XX Best Local Similarity 100.0%; Pred. No. 36;
XX Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 3 ctggcccccacagctcta 20
DB 1 ctggcccccacagctcta 18

RESULT 2
ID AAA62444/C
AC AAA62444 standard; DNA; 25 BP.
AC AAA62444;
DT 13-NOV-2000 (first entry)
DE Human Akt-3 cDNA 3'RACE sense primer Akt-3sp4.
XX Human: Akt-3; protein kinase B; PKB; serine/threonine kinase; cytosolic;
XX apoptosis stimulator; cancer; rapid amplification of cDNA ends; RACE;
XX chromosome 1q43-44; PCR primer; ss.
XX Homo sapiens.
XX WO200037613-A2.
XX 29-JUN-2000.
XX 17-DEC-1999; 99WO-GB04311.
XX 22-DEC-1998; 98GB-0028375.
XX (JANC) JANSSEN PHARM NV.
XX Measure SLJ, Richardson A;
XX WPI; 2000-498840/44.
XX New human serine/threonine kinase protein and the polynucleotide
XX encoding the protein, useful for preparing a medicament for treating
XX disorders associated with human serine/threonine kinase protein
XX activity, especially cancer.

PS Disclosure; Page 20; 61pp; English.

XX The present sequence is a primer used to isolate human Akt-3 cDNA from
CC human brain cDNA by 3' rapid amplification of cDNA ends (3' RACE). Akt-3
CC is a third human isoform of Akt, which is also known as protein kinase B
CC (PKB) or "related to A and C protein kinase" (RAC-PK). It is located on
CC human chromosome 1, region q43-q44. The present primer was used in the
CC second round of 3' RACE. The sequence is based on the product of the
CC first round, which was performed using primers based on a human
CC hippocampal EST sequence that showed high similarity to the rat
CC RAC-PKgamma sequence. Primers based on the product of the second round of
CC 3' RACE were then used to amplify the complete coding sequence of Akt-3

CC from human hippocampal cDNA. Akt can inhibit apoptosis induced by
CC detachment from the extracellular matrix. The Akt-3 nucleic acid molecule
CC and protein may be used as medicaments for treating cancer. Agents which
CC influence the activity of Akt-3 protein, and so stimulate apoptosis, may
CC also be used to treat diseases associated with Akt-3.

XX Sequence 25 BP; 5 A; 5 C; 10 G; 5 T; 0 other;

XX Query Match 64.0%; Score 16; DB 21; Length 25;
XX Best Local Similarity 100.0%; Pred. No. 2.8e+02;
XX Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 10 ccacagctcactgct 25
DB 25 CCACAGCTCCTACTGCT 10

RESULT 3
ID AAQ97960
AC AAQ97960 standard; DNA; 20 BP.
AC AAQ97960;
DT 18-OCT-1995 (first entry)
DE PNA oligomer targeting coding region of PKC-epsilon.
XX Peptide nucleic acid; PNA; PKC-alpha; protein kinase C; ss;
XX cell proliferation; cell differentiation; isozyme; antisense;
XX triple helix; cancer; psoriasis; inflammation.
XX Synthetic.
XX OS

PH Key Location/Qualifiers
FT misc-feature 1..20
FT /*tag= a
FT /note= "at least one (and preferably all) of
FT the backbone subunits are composed of N-acetyl
FT N-(2-aminoethyl)glycine peptide residues, the
FT nucleobase being attached covalently to the
FT acetyl group and the peptide linkage being
FT formed by condensation of the glycine
FT carboxy group of one residue with the amino
FT group of the 2-aminoethyl moiety in the next
FT residue"

XX WO9503833-A.
XX 09-FEB-1995.
XX 28-JUL-1994; 94WO-US08465.
XX 29-JUL-1993; 93US-0099098.
XX (ISIS-) ISIS PHARM INC.
XX Dean NM;
XX WPI; 1995-082040/11.
XX New peptide nucleic acid oligomers specific for protein kinase C
XX isozyme(s) - useful as antisense molecules for treating PKC
XX mediated disease, e.g. cancer, psoriasis and inflammation

PS Claim 38; Page 274; 287pp; English.

XX New peptide nucleic acid (PNA) oligomers are provided which (a) consist
CC of naturally occurring nucleobases covalently bound to a polyamide
CC backbone and (b) hybridise to the translation initiation AUG region,
CC coding region, 5' untranslated region (5' UTR) or 3' untranslated region
CC (3' UTR) of PKC-alpha or its isoforms. The PNAs can be used to target
CC RNA and single stranded DNA (ssDNA) to produce antisense-type gene

CC regulation molecules. They inhibit expression of PKC-alpha and its
CC isoforms (including beta, gamma, delta, epsilon, zeta and eta) and so
CC are useful for treating and diagnosing cell proliferation and
CC differentiation processes such as neoplastic, hyperproliferative
CC and inflammatory diseases.
CC PNA oligomers have high affinity for complementary single stranded DNA.
CC They are also able to form triple helices in which a first PNA strand
CC binds with RNA or ssDNA and a second PNA strand binds with the resulting
CC double helix or with the first PNA strand. The PNAs possess no
CC significant charge and are water soluble, which facilitates cellular
CC uptake. Further, since they contain amides of non-biological amino acids,
CC they are biostable and resistant to enzymatic degradation by proteases.
CC The present sequence targets the coding region of PKC-epsilon.
CC
XX
SQ Sequence 20 BP; 4 A; 11 C; 4 G; 1 T; 0 other;

Query Match 60.8%; Score 15.2; DB 16; Length 20;
Best Local Similarity 85.0%; Pred. No. 6.2e+02;
Matches 17; Conservative 0; Mismatches 3; Indels 0; Gaps 0;
QY 1 ccctagggcccccagctca 20
||| ||| ||||| ||||| ||
DB 1 ccccgagggcccccagctcca 20

RESULT 4

AA084237 standard; DNA; 20 BP.

AA084237;

21-SEP-1995 (first entry)

PKC-epsilon coding region antisense oligo, ISIS #7944.

Antisense; protein kinase C; alpha; PKC; beta; gamma; eta; epsilon;

zeta; modulation; expression; isozyme; hybridise; 5' UTR; human;

3' untranslated region; translation initiation site; detection;

phosphorothioate linkage; 2'-O-methyl modification;

2'-O-propyl modification; ss.

Synthetic.

WO9502069-A.

19-JAN-1995.

08-JUL-1994; 94WO-US07770.

09-JUL-1993; 93US-0089996.

22-FEB-1994; 94US-019779.

(ISIS-) ISIS PHARM INC.

Bennett CF, Boggs RT, Dean NM;

WPI; 1995-066911/09.

Oligo:nucleotide(s) hybridisable with Protein Kinase C mRNA or

gene - also novel PKC-alpha 3'-UTR sequence, useful for

diagnosis and treatment of hyperproliferative disorders.

Claim 115; Page 37; 125pp; English.

The sequences given in AA084236-40 are oligos which are antisense to the

protein kinase C-epsilon (PKC-epsilon) cDNA. These antisense molecules

may be used in modulating the expression of this particular isozyme of

PKC. The oligos of the invention preferably hybridise with the 5'- or

3'-untranslated regions of the PKC gene, or the translation initiation

site, or the coding region. These oligos may be used in the detection

of the human PKC genes and for treatment of animals with conditions

associated with PKC, esp. hyperproliferative diseases such as psoriasis,

CC colorectal cancer, lung cancer, breast or skin cancer. These oligos may
CC contain at least one phosphorothioate linkage and/or at least one of the
CC nucleotides comprises a modification on the 2' position of the sugar,
CC esp. a 2'-O-methyl or a 2'-O-propyl modification.
CC
XX
SQ Sequence 20 BP; 4 A; 11 C; 4 G; 1 T; 0 other;

Query Match 60.8%; Score 15.2; DB 16; Length 20;
Best Local Similarity 85.0%; Pred. No. 6.2e+02;
Matches 17; Conservative 0; Mismatches 3; Indels 0; Gaps 0;
QY 1 ccctagggcccccagctca 20
||| ||| ||||| ||||| ||
DB 1 ccccgagggcccccagctcca 20

RESULT 5

AA227354 standard; DNA; 20 BP.

AA227354;

01-DEC-1999 (first entry)

Human protein kinase C epsilon antisense oligonucleotide #12.

Human; protein kinase C; PKC; diagnosis; antisense oligonucleotide;

phosphorothioate; hybridisation; isozyme; target; inflammation;

hyperproliferative disorder; psoriasis; tumour; cancer; glioblastoma; ss.

Synthetic.

Homo sapiens.

US5959096-A.

28-SEP-1999.

07-JUN-1995; 95US-0481066.

16-MAR-1992; 92US-0852852.

09-JUL-1993; 93US-0089996.

(ISIS-) ISIS PHARM INC.

Bennett CF, Dean N;

WPI; 1999-561076/47.

Example 16; Column 23; 56pp; English.

The present invention describes antisense oligonucleotides up to 50

nucleotides in length which specifically bind mRNA encoding human

protein kinase C (PKC). AA227266 to AA227386 represent human PKC

antisense oligonucleotides used in the exemplification of the present

invention. The antisense oligonucleotides are useful for the treatment of

diseases associated with PKC expression, such as hyperproliferative and

inflammatory conditions including psoriasis, tumours and cancer

(glioblastoma, bladder, breast, colon and lung cancer).

Sequence 20 BP; 4 A; 11 C; 4 G; 1 T; 0 other;

Query Match 60.8%; Score 15.2; DB 20; Length 20;
Best Local Similarity 85.0%; Pred. No. 6.2e+02;
Matches 17; Conservative 0; Mismatches 3; Indels 0; Gaps 0;
QY 1 ccctagggcccccagctca 20
||| ||| ||||| ||||| ||
DB 1 ccccgagggcccccagctcca 20

RESULT 6
AAx78612
ID AAX78612 standard; DNA: 20 BP.
XX
AC AAX78612:
XX
DT 03-SEP-1999 (first entry)
XX
DE Human PKC-epsilon oligonucleotide primer ISIS # 7944.
XX
KW PKC; human; PKC-alpha; primer; protein kinase C; expression modulator;
KW PKC-beta type I; PKC-beta type II; PKC-gamma; PKC-eta; PKC-delta;
KW PKC-epsilon; PKC-zeta; anti-inflammatory; cytosolic; antisense targeting;
KW isozyme; growth control; hyperproliferative disease; colon cancer;
KW glioblastoma; bladder cancer; inflammatory condition; psoriasis; ss.
XX
OS Synthetic.
OS Homo sapiens.
XX
PN US5922686-A.
XX
PD 13-JUL-1999.
XX
PF 14-JUN-1996; 96US-0664336.
XX
PR 14-JUN-1996; 96US-0664336.
PR 16-MAR-1992; 92US-0852852.
PR 09-JUL-1993; 93US-0089996.
XX
PA (ISIS-) ISIS PHARM INC.
XX
PI Bennett CF, Dean N;
XX
DR WPI; 1999-404471/34.
XX
PT Oligonucleotides targeted against nucleic acids encoding protein
PT kinase C
XX
PS Example 16; Column 63-64; 56pp; English.
XX
CC This invention describes novel oligonucleotides (AAX78524-X78644) having
CC up to 50 nucleotides hybridisable with, and able to modulate the
CC expression of, a nucleic acid encoding protein kinase C and its isozymes
CC alpha, beta type I, beta type II, gamma, eta, delta, epsilon and zeta.
CC The oligonucleotides of the invention have anti-inflammatory and
CC cytostatic activity and are used for antisense targeting to modulate the
CC expression of PKC or of a particular PKC isozyme or set of isozymes in
CC cells or tissues. The products of the invention also hybridise with
CC nucleic acids involved in the modulation of PKC expression, which is
CC known to be involved in growth control in hyperproliferative diseases e.g.
CC colon cancer, glioblastoma and bladder cancer as well as in inflammatory
CC conditions e.g. psoriasis. Due to their specificity the oligonucleotides
CC are able to overcome the problems of toxicity associated with previous
CC agents designed to modulate PKC expression.
XX
SQ Sequence 20 BP; 4 A; 11 C; 4 G; 1 T; 0 other;

Query Match 60.8%; Score 15.2; DB 20; Length 20;
Best Local Similarity 85.0%; Pred. No. 6.2e+02;
Matches 17; Conservative 0; Mismatches 3; Indels 0; Gaps 0;

OY 1 cccatggcccccagctca 20
||| ||| ||| ||| ||| ||| |||
Db 1 ccccaaggcccccagctca 20

RESULT 7
AAX83704
ID AAX83704 standard; DNA: 20 BP.
XX

AC AAX83704;
XX
DT 27-AUG-1999 (first entry)
XX
DE Human protein kinase C antisense oligonucleotide SEQ ID NO:89.
XX
KW Human; protein kinase C; PKC; antisense oligonucleotide; diagnosis; ss;
KW hybridisation; cancer; psoriasis; hyperproliferative disease; tumour.
XX
OS Synthetic.
OS Homo sapiens.
XX
PN US5916807-A.
XX
PD 29-JUN-1999.
XX
PF 07-JUN-1995; 95US-0481072.
XX
PR 07-JUN-1995; 95US-0481072.
PR 16-MAR-1992; 92US-0852852.
PR 09-JUL-1993; 93US-0089996.
XX
PA (ISIS-) ISIS PHARM INC.
XX
PI Bennett CF, Dean N;
XX
DR WPI; 1999-403817/34.
XX
PT New antisense oligonucleotides specific for human protein kinase C
PT useful for diagnosis and treatment of cancer and psoriasis
XX
PS Claim 1; Column 21; 54pp; English.
XX
CC The present invention describes a method of inhibiting the expression of
CC human protein kinase C (PKC) in cells. The method comprises contacting
CC the cells with an antisense oligonucleotide which has up to 50
CC nucleotide units. AAX83633 to AAX83720 represent specifically claimed
CC antisense oligonucleotides for use in the method of the invention.
CC The antisense oligonucleotides modulate hybridize to messenger RNA from
CC the PKC gene which results in modulation of expression of the PKC gene.
CC This means they can be used for diagnosis, therapeutic or prophylactic
CC treatment of PKC associated diseases such as cancer and psoriasis, and
CC as research agents. Abnormal proliferative states in tissue from
CC patients suspected of having a hyperproliferative disease e.g. cancer,
CC psoriasis can be diagnosed. Tumours associated with PKC can be
CC distinguished from tumours which are not PKC associated to allow an
CC efficacious treatment regime to be used. The antisense oligonucleotides
CC have specific activity so are able to modulate PKC activity without
CC producing side effects and with greater effectiveness than observed
CC from administration of current agents. AAX83721 to AAX83753 represent
CC other oligonucleotides used in examples from the present invention.
XX
SQ Sequence 20 BP; 4 A; 11 C; 4 G; 1 T; 0 other;

Query Match 60.8%; Score 15.2; DB 20; Length 20;
Best Local Similarity 85.0%; Pred. No. 6.2e+02;
Matches 17; Conservative 0; Mismatches 3; Indels 0; Gaps 0;

OY 1 cccatggcccccagctca 20
||| ||| ||| ||| ||| ||| |||
Db 1 ccccaaggcccccagctca 20

RESULT 8
AAX22650
ID AAX22650 standard; DNA: 20 BP.
XX
AC AAX22650;
XX
DT 27-MAY-1999 (first entry)
XX
DE Human protein kinase C antisense oligonucleotide #89.

XX Protein kinase C; PKC; human; antisense; primer; inhibitor; treatment;
 KW hyperproliferative condition; cancer; colorectal; breast; bladder; lung;
 KW brain; glioblastoma multiforme; skin; psoriasis; ss.
 XX Synthetic.
 OS Homo sapiens.
 XX US585970-A.
 PN 23-MAR-1999.
 XX 07-JUN-1995; 95US-0488177.
 XX 07-JUN-1995; 95US-0488177.
 PR 16-MAR-1992; 92US-0852852.
 PR 09-JUL-1993; 93US-0089996.
 XX (ISIS-) ISIS PHARM INC.
 PA Bennett CF, Dean N;
 PI WPI, 1999-228583/19.
 DR WPI, 1999-228583/19.
 XX New human protein kinase C antisense oligonucleotides - useful for
 PT treating PKC-related hyperproliferative conditions e.g. cancer and
 PS psoriasis
 XX Example 16; Column 21; 55pp; English.
 XX This invention describes antisense oligonucleotides that specifically
 CC bind to human protein kinase C (PKC) mRNA. These oligonucleotides can be
 CC used to inhibit PKC mRNA and therefore be used to treat PKC-related
 CC hyperproliferative conditions, e.g. cancer, especially colorectal cancer,
 CC breast cancer, bladder cancer, lung cancer, or brain cancer (preferably
 CC glioblastoma multiforme). The products of the invention may also be used
 CC to treat skin cancer and psoriasis.
 XX Sequence 20 BP; 4 A; 11 C; 4 G; 1 T; 0 other:
 SQ

Query Match 60.8%; Score 15.2; DB 20; Length 20;
 Best Local Similarity 85.0%; Pred. No. 6.2e+02;
 Matches 17; Conservative 0; Mismatches 3; Indels 0; Gaps 0;

QY 1 cccctagggccaccagctca 20
 ||| ||| ||| ||| ||| ||| |||
 Db 1 ccccgagggccaccagctcca 20

RESULT 9
 AAX19215
 ID AAX19215 standard; DNA; 20 BP.
 AC AAX19215;
 XX 14-MAY-1999 (first entry)
 DT Human PKC-epsilon antisense oligonucleotide SEQ ID NO:89.
 XX Human PKC-epsilon antisense oligonucleotide;
 DE Human PKC-epsilon antisense oligonucleotide;
 KW Human PKC-epsilon antisense oligonucleotide;
 KW phosphocholate linkage; hyperproliferative disease; cancer;
 KW psoriasis; tumour; inhibition; ss.
 XX Synthetic.
 OS Homo sapiens.
 XX US5882927-A.
 PN 16-MAR-1999.
 PD 16-MAR-1999.
 PF 08-JUN-1995; 95US-0478178.
 XX

PR 07-JUN-1995; 95US-0478178.
 PR 16-MAR-1992; 92US-0852852.
 PR 09-JUL-1993; 93US-0089996.
 XX (ISIS-) ISIS PHARM INC.
 PA Bennett CF, Dean N;
 PI WPI, 1999-214073/18.
 DR WPI, 1999-214073/18.
 XX New synthetic oligonucleotides inhibiting expression of protein
 PT kinase C (PKC)-alpha - useful for treating and diagnosing conditions
 PS associated with abnormal PKC expression
 XX Example 16; Column 23; 56pp; English.
 XX The present invention specifically describes antisense oligonucleotides
 CC of up to 50 nucleotides in length which specifically bind human protein
 CC kinase C-alpha (PKC-alpha) mRNA. AAX19127 to AAX19247 represent
 CC antisense oligonucleotides from the present invention which bind human
 CC PKC-alpha, -delta, -gamma, -epsilon, -zeta and -eta. The
 CC antisense oligonucleotides modulate the expression of the PKC gene (i.e.
 CC inhibit the PKC gene). The antisense oligonucleotides can be used to
 CC diagnose abnormal proliferative states in tissue or other samples from
 CC patients suspected of having a hyperproliferative disease e.g. cancer or
 CC psoriasis. The antisense oligonucleotides can be used to distinguish
 CC PKC-associated tumours and to detect and diagnose PKC expression (through
 CC the use of 32P labeled antisense oligonucleotides). Radiolabeled
 CC antisense oligonucleotides can also be used to perform autoradiography of
 CC tissues to determine the localization, distribution and quantitation of
 CC PKC expression for research, diagnostic and therapeutic purposes. The use
 CC of the antisense oligonucleotides eliminate the side effects associated
 CC with prior art methods because it modulates the amount of PKC protein
 CC made from the gene rather than inhibiting the enzyme itself.
 XX Sequence 20 BP; 4 A; 11 C; 4 G; 1 T; 0 other:
 SQ

Query Match 60.8%; Score 15.2; DB 20; Length 20;
 Best Local Similarity 85.0%; Pred. No. 6.2e+02;
 Matches 17; Conservative 0; Mismatches 3; Indels 0; Gaps 0;

QY 1 cccctagggccaccagctca 20
 ||| ||| ||| ||| ||| ||| |||
 Db 1 ccccgagggccaccagctcca 20

RESULT 10
 AAT53201/C
 ID AAT53201 standard; RNA; 36 BP.
 AC AAT53201;
 XX 02-MAY-1997 (first entry)
 DT Mouse ICAM hammerhead ribozyme sequence (nt. position 2378).
 XX Enzymatic nucleic acid; ribozyme; trans cleavage; inhibition;
 KW gene expression; downregulation; interleukin-5; IL-5; ICAM-1;
 KW intercellular adhesion molecule; rel A; tumour necrosis factor;
 KW TNF-alpha; respiratory syncytial virus; RSV; bcr-abl; oncogene;
 KW translocation; chronic myelogenous leukaemia; CML; cancer;
 KW Philadelphia chromosome; inflammation; autoimmune disease;
 KW atherosclerosis; myocardial infarction; stroke; restenosis;
 KW transplant rejection; rheumatoid arthritis; psoriasis;
 KW myocardial ischaemia; Kawasaki disease; septic shock; HIV;
 KW human immunodeficiency virus; acquired immune deficiency syndrome;
 KW AIDS; ss.
 XX Synthetic.
 OS Synthetic.
 XX WO9523225-A2.
 PN

PD		31-AUG-1995.	
XX			
PF	23-FEB-1995;	95WO-IB00156.	
XX			
PR	30-JAN-1995;	95US-0380734.	
PR	23-FEB-1994;	94US-0201109.	
PR	29-MAR-1994;	94US-0218934.	
PR	04-APR-1994;	94US-0222795.	
PR	15-APR-1994;	94US-0224483.	
PR	15-APR-1994;	94US-0227958.	
PR	15-APR-1994;	94US-0228041.	
PR	18-MAY-1994;	94US-0245736.	
PR	06-JUL-1994;	94US-0271280.	
PR	15-AUG-1994;	94US-0291932.	
PR	16-AUG-1994;	94US-0291433.	
PR	17-AUG-1994;	94US-0292620.	
PR	19-AUG-1994;	94US-0293520.	
PR	08-SEP-1994;	94US-0300000.	
PR	08-SEP-1994;	94US-0303039.	
PR	23-SEP-1994;	94US-0311466.	
PR	23-SEP-1994;	94US-0311749.	
PR	28-SEP-1994;	94US-0314397.	
PR	03-OCT-1994;	94US-0316771.	
PR	07-OCT-1994;	94US-0319492.	
PR	11-OCT-1994;	94US-0321993.	
PR	04-NOV-1994;	94US-0334847.	
PR	10-NOV-1994;	94US-0337608.	
PR	28-NOV-1994;	94US-0345516.	
PR	16-DEC-1994;	94US-0357577.	
PR	23-DEC-1994;	94US-0363233.	
PA	(RIBO-) RIBOZYME PHARM INC.		
XX			
PI	Stinchcomb DT, Chowrira B, Direnzo A, Draper KG, Dutycz LW;		
PI	Grimm S, Karpelesky A, Kisch K, Matulich-Adamic J;		
PI	Mcsivigen JA, Modak A, Pavco P, Beigelman L, Sullivan SM;		
PI	Swedler D, Thompson JD, Tracz D, Usman N, Wincott FE;		
PI	Woolf T;		
XX			
DR	WPI: 1995-351090/45.		
XX			
PT	Ribozymes having modified bases and methods for producing them -		
PT	for use in inhibiting disease related genes		
XX			
PS	Claim 9; Page 195; 407pp; English.		
XX			
CC	The present sequence is that of a claimed enzymatic nucleic acid		
CC	(i.e. a ribozyme) which cleaves ICAM-1 mRNA at the nucleotide base		
CC	position indicated in the DE line.		
CC	Regions of the mRNA that do not form secondary folding		
CC	structures and that contain potential hammerhead and hairpin		
CC	ribozyme cleavage sites were identified by computer analysis.		
CC	Ribozymes directed against these mRNA sequences were designed and		
CC	synthesised with modifications that improve their nuclease		
CC	resistance. The ribozymes cleave the ICAM-1 target sequences and		
CC	thereby inhibit ICAM-1 expression, making them useful for reducing		
CC	transplant rejection and alleviating symptoms in patients with		
CC	rheumatoid arthritis, asthma and other inflammatory disorders.		
XX			
SO	Sequence 36 BP; 13 A; 9 C; 10 G; 4 U; 0 other;		
OY	Query Match	60.8%; Score 15.2; DB 16; Length 36;	
	Best Local Similarity	85.0%; Pred. No. 6.4e+02;	
DB	Matches 17; Conservative 0; Mismatches 3; Indels 0; Gaps 0.		
	4 tagcgcaccacgcttactg 23		
	20 TCggccTATCAGTCGTACTG 1		

RESULT 11
AA#52969/C

ID	AA52969 standard; RNA; 36 Bc.
XX	
AC	AA52969;
XX	
DT	21-APR-1997 (first entry)
XX	
XX	Mouse ICAM hammerhead ribozyme sequence (nt. position 263).
XX	
KW	Enzymatic nucleic acid; ribozyme; trans cleavage; inhibition;
KW	gene expression; downregulation; interleukin-5; IL-5; ICAM-1;
KW	intercellular adhesion molecule; rel A; tumor necrosis factor;
KW	TNF-alpha; respiratory syncytial virus; RSV; bcr-abl; oncogene;
KW	translocation; chronic myelogenous leukaemia; CML; cancer;
KW	atherosclerosis; myocardial infarction; stroke; restenosis;
KW	transplant rejection; rheumatoid arthritis; psoriasis;
KW	myocardial ischaemia; Kawasaki disease; septic shock; HIV;
KW	human immunodeficiency virus; acquired immune deficiency syndrome;
AD	AIDS; ss.
XX	
OS	Synthetic.
XX	
PN	W09523225-A2.
XX	
PF	31-AUG-1995.
XX	
PF	23-FEB-1995; 95WO-1B00156.
XX	
PR	30-JAN-1995; 95US-0380734.
PR	23-FEB-1994; 94US-0201109.
PR	29-MAR-1994; 94US-0218934.
PR	04-APR-1994; 94US-0222795.
PR	07-APR-1994; 94US-0224483.
PR	15-APR-1994; 94US-0227958.
PR	15-APR-1994; 94US-0228041.
PR	18-MAY-1994; 94US-0245736.
PR	06-JUL-1994; 94US-0211280.
PR	15-AUG-1994; 94US-0291932.
PR	16-AUG-1994; 94US-0291433.
PR	17-AUG-1994; 94US-0293520.
PR	19-AUG-1994; 94US-0293520.
PR	02-SEP-1994; 94US-0300000.
PR	08-SEP-1994; 94US-0303039.
PR	23-SEP-1994; 94US-0310456.
PR	28-SEP-1994; 94US-0311749.
PR	28-SEP-1994; 94US-0314397.
PR	03-OCT-1994; 94US-0316771.
PR	07-OCT-1994; 94US-0319492.
PR	11-OCT-1994; 94US-0321993.
PR	10-NOV-1994; 94US-0334847.
PR	10-NOV-1994; 94US-0337608.
PR	28-NOV-1994; 94US-0345516.
PR	16-DEC-1994; 94US-0357577.
PR	23-DEC-1994; 94US-0363233.
XX	
PA	(RIBO-) RIBOZYME PHARM INC.
XX	
PI	Stinchcomb DT, Chowrita B, Dizenzo A, Draper KG, Dudycz LW;
PI	Grimm S, Kappelesky A, Ksich K, Matulis-Adamic J;
PI	McGwigen JA, Modak A, Payco P, Belgelman L, Sullivan SM;
PI	Swedder D, Thompson JD, Tracz D, Usman N, Wincott FE;
PI	Woolf T;
XX	
DR	WPI; 1995-351090/45.
XX	
PT	Ribozymes having modified bases and methods for producing them
PT	for use in inhibiting disease related genes
XX	
PS	Claim 9; Page 190; 407pp; English.
XX	
CC	The present sequence is that of a claimed enzymatic nucleic acid
CC	(i.e. a ribozyme) which cleaves ICAM-1 mRNA at the nucleotide base
CC	position indicated in the DE line.

SQ Sequence 51 BP; 12 A; 16 C; 14 G; 9 T; 0 other;

Query Match 60.0%; Score 15; DB 22; Length 51;
Best Local Similarity 78.3%; Pred. No. 8e+02;
Matches 18; Conservative 0; Mismatches 5; Indels 0; Gaps 0;

QY 2 cctagagcccccacagctctactgc 24
111 111 111 111 111 111
DB 28 CCGTGGCCCTACCAGGCTGCTGC 6

RESULT 14

AAZ27368
ID AAZ27368 standard; DNA; 20 BP.

AC AAZ27368;

DT 01-DEC-1999 (first entry)

DE Human protein kinase C epsilon antisense oligonucleotide #26.

KW Human: protein kinase C; PKC; diagnosis; antisense oligonucleotide;

KM phosphotriester; hybridisation; isozyme; target; inflammation;
hyperproliferative disorder; psoriasis; tumour; cancer; glioblastoma; ss.

OS Synthetic.

OS Homo sapiens.

PN US9599096-A.

PD 28-SEP-1999.

PF 07-JUN-1995; 95US-0481066.

PR 16-MAR-1992; 92US-0852852.

PR 09-JUL-1993; 93US-0089996.

XX (ISIS-) ISIS PHARM INC.

PI Bennett CF, Dean N;

DR WPI: 1999-561076/47.

PT Antisense oligonucleotides useful for treatment of hyperproliferative

PT and inflammatory conditions including psoriasis, tumours and cancer -

XX Example 16; Column 23; 56pp; English.

XX The present invention describes antisense oligonucleotides up to 50

CC nucleotides in length which specifically bind mRNA encoding human

CC protein kinase C (PKC). AAZ27266 to AAZ27386 represent human PKC

CC antisense oligonucleotides used in the exemplification of the present

CC invention. The antisense oligonucleotides are useful for the treatment of

CC diseases associated with PKC expression, such as hyperproliferative and

CC inflammatory conditions including psoriasis, tumours and cancer

CC (glioblastoma, bladder, breast, colon and lung cancer).

RESULT 15

AAZ78626
ID AAZ78626 standard; DNA; 20 BP.

XX AAX78626;

DT 03-SEP-1999 (first entry)

DE Human PKC-epsilon oligonucleotide primer ISIS H.

KW PKC: human; PKC-alpha; primer; protein kinase C; expression modulator;

KM PKC-beta type I; PKC-beta type II; PKC-gamma; PKC-eta; PKC-delta;

KM PKC-epsilon; PKC-zeta; anti-inflammatory; cytoskeletal; antisense targeting;

KM isozyme; growth control; hyperproliferative disease; colon cancer;

KM glioblastoma; bladder cancer; inflammatory condition; psoriasis; ss.

OS Synthetic.

OS Homo sapiens.

PN US922686-A.

PD 13-JUL-1999.

PF 14-JUN-1996; 96US-0664336.

PR 14-JUN-1996; 96US-0664336.

PR 16-MAR-1992; 92US-0852852.

PR 09-JUL-1993; 93US-0089996.

XX (ISIS-) ISIS PHARM INC.

PI Bennett CF, Dean N;

DR WPI: 1999-404471/34.

PT Oligonucleotides targeted against nucleic acids encoding protein

PT kinase C

XX Example 16; Column 69-70; 56pp; English.

XX This invention describes novel oligonucleotides (AAZ78524-X78644) having

CC up to 50 nucleotides hybridisable with, and able to modulate the

CC expression of, a nucleic acid encoding protein kinase C and its isozymes

CC alpha, beta type I, beta type II, gamma, eta, delta, epsilon and zeta.

CC The oligonucleotides of the invention have anti-inflammatory and

CC cytoskeletal activity and are used for antisense targeting to modulate the

CC expression of PKC or of a particular PKC isozyme or set of isozymes in

CC cells or tissues. The products of the invention also hybridise with

CC nucleic acids involved in the modulation of PKC expression, which is

CC known to be involved growth control in hyperproliferative diseases e.g.

CC colon cancer, glioblastoma and bladder cancer as well as in inflammatory

CC conditions e.g. psoriasis. Due to their specificity the oligonucleotides

CC are able to overcome the problems of toxicity associated with previous

XX agents designed to modulate PKC expression.

SQ Sequence 20 BP; 3 A; 12 C; 4 G; 1 T; 0 other;

Query Match 59.2%; Score 14.8; DB 20; Length 20;
Best Local Similarity 88.9%; Pred. No. 9.4e+02;
Matches 16; Conservative 0; Mismatches 2; Indels 0; Gaps 0;

QY 1 ccttaggcccacacagtc 18
111 111 111 111 111 111
DB 2 ccccgagggccacacagtc 19

Search completed: March 9, 2002, 01:06:58
Job time: 11944 sec

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